

MAGLUMI[®] MPO (CLIA)

INTENDED USE

The kit is an *in vitro* chemiluminescence immunoassay for the quantitative determination of Myeloperoxidase (MPO) in human plasma using the MAGLUMI series Fully-auto chemiluminescence immunoassay analyzer and Biolumi series Integrated System, and the assay is used for an aid in the diagnosis of cardiovascular inflammation.

SUMMARY

Myeloperoxidase (MPO) is a heme-containing protease secreted by neutrophils, monocytes and certain tissue macrophages. It is a member of the mammalian heme peroxidase superfamily ^{1, 2}. MPO is a tetramer composed of two glycosylated 59- to 64- kDa heavy chain and two nonglycosylated 14-kDa light chains, and the mature MPO molecule has a molecular mass of approximately 150 kDa ^{3, 4}. Activation of leukocytes promotes the release of myeloperoxidase (MPO) from leukocyte granules, catalyzing the production of reactive oxygen species and oxygen free radicals ⁵. MPO is one of several proteins or enzymes which show antimicrobial properties in neutrophils.

Inflammatory events have been implicated at all stages in the evolution of atherosclerotic plaque ⁶. Cardiovascular inflammation may evolve into atherosclerosis, coronary artery disease (CAD), heart failure and stroke. CAD is one of the major cardiovascular diseases affecting the global human population. CAD is an atherosclerotic disease which is inflammatory in nature, manifested by stable angina, unstable angina, myocardial infarction (MI), or sudden cardiac death ⁷. Elevated level of myeloperoxidase has a significant correlation with CAD; CAD patients had significantly higher concentrations of MPO compared to the controls ^{8,9}. MPO participates in the progression of atherosclerosis and rupture the atherosclerotic plaques, leading to elevated MPO levels in acute coronary syndrome (ACS) patients ¹⁰. In patients with ACS, MPO serum levels powerfully predict an increased risk for subsequent cardiovascular events. MPO may serve as both a marker and mediator of vascular inflammation and further points toward the significance of the pathophysiology of ACS.

TEST PRINCIPLE

Sandwich chemiluminescence immunoassay.

The sample, magnetic microbeads coated with anti-MPO antibody, ABEI labeled with another anti-MPO antibody and buffer are mixed thoroughly and incubated, forming sandwich complexes. After precipitation in a magnetic field, the supernatant is decanted and then perform a wash cycle. Subsequently, the Starter 1+2 are added to initiate a chemiluminescent reaction. The light signal is measured by a photomultiplier as relative light units (RLUs), which is proportional to the concentration of MPO present in the sample.

REAGENTS

Kit Contents

Component	Description	100 tests/kit	50 tests/kit	30 tests/kit	
Magnetic Microbeads	Magnetic microbeads coated with anti-MPO antibody (~8.00 μ g/mL) in PBS buffer, NaN ₃ (<0.1%).	2.5 mL	2.0 mL	1.0 mL	
Calibrator Low	A low concentration of MPO antigen in PBS buffer, NaN_3 (<0.1%).	1.0 mL	1.0 mL	1.0 mL	
Calibrator High	A high concentration of MPO antigen in PBS buffer, NaN ₃ (<0.1%).	1.0 mL	1.0 mL	1.0 mL	
Buffer	PBS buffer, NaN ₃ (<0.1%).	13.5 mL	7.5 mL	3.0 mL	
ABEI Label	ABEI labeled with anti-MPO antibody (~18.8 ng/mL) in Tris-HCI buffer, NaN ₃ (<0.1%).	23.5 mL	12.5 mL	6.3 mL	
Diluent	PBS buffer, NaN ₃ (<0.1%).	15.0 mL	10.0 mL	3.5 mL	
Control 1	A low concentration of MPO antigen (150 ng/mL) in PBS buffer, NaN ₃ (<0.1%).	1.0 mL	1.0 mL	1.0 mL	
Control 2	A high concentration of MPO antigen (600 ng/mL) in PBS buffer, NaN ₃ (<0.1%).	1.0 mL	1.0 mL	1.0 mL	
All reagents are provided ready-to-use					

Warnings and Precautions

- For *in vitro* diagnostic use.
- For professional use only.
- Exercise the normal precautions required for handling all laboratory reagents.
- Personal protective measures should be taken to prevent any part of the human body from contacting samples, reagents, and controls, and should comply with local operating requirements for the assay.
- · A skillful technique and strict adherence to the package insert are necessary to obtain reliable results.
- Do not use kit beyond the expiration date indicated on the label.
- · Do not interchange reagent components from different reagents or lots.
- Avoid foam formation in all reagents and sample types (specimens, calibrators and controls).
- All waste associated with biological samples, biological reagents and disposable materials used for the assay should be considered potentially infectious and should be disposed of in accordance with local guidelines.
- This product contains sodium azide. Sodium azide may react with lead or copper plumbing to form highly explosive metal azides. Immediately after disposal, flush
 with a large volume of water to prevent azide build-up. For additional information, see Safety Data Sheets available for professional user on request.

Note: If any serious incident has occurred in relation to the device, please report to Shenzhen New Industries Biomedical Engineering Co., Ltd. (Snibe) or our authorized representative and the competent authority of the Member State in which you are established.

Reagent Handling

- To avoid contamination, wear clean gloves when operating with a reagent kit and sample. When handling reagent kit, replace the gloves that have been in contact
 with samples, since introduction of samples will result in unreliable results.
- Do not use kit in malfunction conditions; e.g., the kit leaking at the sealing film or elsewhere, obviously turbid or precipitation is found in reagents (except for Magnetic Microbeads) or control value is out of the specified range repeatedly. When kit in malfunction conditions, please contact Snibe or our authorized distributor.
- To avoid evaporation of the liquid in the opened reagent kits in refrigerator, it is recommended that the opened reagent kits to be sealed with reagent seals contained within the packaging. The reagent seals are single use, and if more seals are needed, please contact Snibe or our authorized distributor.
- Over time, residual liquids may dry on the septum surface. These are typically dried salts and have no effect on assay efficacy.
- Use always the same analyzer for an opened reagent integral.
- For magnetic microbeads mixing instructions, refer to the Preparation of the Reagent section of this package insert.
- · For further information about the reagent handing during system operation, please refer to Analyzer Operating Instructions.

Storage and Stability

- Do not freeze the integral reagents.
- Store the reagent kit upright to ensure complete availability of the magnetic microbeads.
- Protect from direct sunlight.

Stability of the Reagents			
Unopened at 2-8°C	until the stated expiration date		
Opened at 2-8°C	6 weeks		

On-board	4 weeks
Stability of Controls	
Unopened at 2-8°C	until the stated expiration date
Opened at 15-25°C	6 hours
Opened at 2-8°C	6 weeks
Frozen at -20°C	3 months
Frozen and thawed cycles	no more than 3 times

SPECIMEN COLLECTION AND PREPARATION

Specimen Types

Only the specimens listed below were tested and found acceptable.

Specimen Types	Collection Tubes		
Plasma	K2-EDTA, Na2-EDTA		
The second time lists down to stand with a selection of second collection to be the time second collection of the time of time of the time of the time of the time of time of the time of			

The sample types listed were tested with a selection of sample collection tubes that were commercially available at the time of testing, i.e. not all available tubes of
all manufacturers were tested. Sample collection systems from various manufacturers may contain differing materials which could affect the test results in some
cases. Follow tube manufacturers' instructions carefully when using collection tubes.

Other sample types are not allowed to use, serum or heparin samples will have significantly higher MPO concentrations¹¹.

Specimen Conditions

- Do not use heat-inactivated samples or grossly hemolyzed/hyperlipidaemia specimens and specimens with obvious microbial contamination.
- Samples must be free of fibrin and other particulate matter.

• To prevent cross contamination, use of disposable pipettes or pipette tips is recommended.

Preparation for Analysis

- Inspect all specimens for foam. Remove foam with an applicator stick before analysis. Use a new applicator stick for each specimen to prevent cross contamination.
- Frozen specimens must be completely thawed before mixing. Mix thawed specimens thoroughly by low speed vortexing or by gently inverting. Visually inspect
 the specimens. If layering or stratification is observed, mix until specimens are visibly homogeneous. If specimens are not mixed thoroughly, inconsistent results
 may be obtained.
- Specimens should be free of fibrin, red blood cells, or other particulate matter. Such specimens may give reliable results and must be centrifuged prior to testing. Transfer clarified specimen to a sample cup or secondary tube for testing. For centrifuged specimens with a lipid layer, transfer only the clarified specimen and not the lipemic material.
- The sample volume required for a single determination of this assay is 10 μL.

Specimen Storage

Specimens removed from the separator, red blood cells or clot may be stored up to 6 hours at 15-25°C or 6 days at 2-8°C, or 6 months frozen at -20°C or colder. Frozen specimens subjected to up to 3 freeze/thaw cycles have been evaluated.

Specimen Shipping

- Package and label specimens in compliance with applicable local regulations covering the transport of clinical specimens and infectious substances.
- Do not exceed the storage limitations listed above.

Specimen Dilution

- Samples, MPO concentrations above the analytical measuring interval, can be diluted with Diluent either automated dilution protocol or manual dilution procedure. The recommended dilution ratio is 1:4. The concentration of the diluted sample must be >260 ng/mL.
- For manual dilution, multiply the result by the dilution factor. For dilution by the analyzers, the analyzer software automatically takes the dilution into account when
 calculating the sample concentration.

PROCEDURE

Materials Provided

MPO (CLIA) assay, control barcode labels.

- Materials Required (But Not Provided)
- General laboratory equipment.
- Fully-auto chemiluminescence immunoassay analyzer Maglumi 600, Maglumi 800, Maglumi 1000, Maglumi 2000, Maglumi 2000 Plus, Maglumi 4000, Maglumi 4000 Plus, MAGLUMI X8, MAGLUMI X3, or Integrated System Biolumi 8000.
- Additional accessories of test required for the above analyzers include Reaction Module, Starter 1+2, Wash Concentrate, Light Check, Tip, and Reaction Cup.
 Specific accessories and accessories' specification for each model refer to corresponding Analyzer Operating Instructions.
- Please use accessories specified by Snibe to ensure the reliability of the test results.

Assay Procedure

- Preparation of the Reagent
- Take the reagent kit out of the box and visually inspect the integral vials for leaking at the sealing film or elsewhere. If there is no leakage, please tear off the sealing film carefully.
- Open the reagent area door; hold the reagent handle to get the RFID label close to the RFID reader (for about 2s); the buzzer will beep; one beep sound indicates
 successful sensing.
- Keeping the reagent straight insert to the bottom along the blank reagent track.
- Observe whether the reagent information is displayed successfully in the software interface, otherwise repeat the above two steps.
- Resuspension of the magnetic microbeads takes place automatically when the kit is loaded successfully, ensuring the magnetic microbeads are totally resuspended homogenous prior to use.

Assay Calibration

- Select the assay to be calibrated and execute calibration operation in reagent area interface. For specific information on ordering calibrations, refer to the calibration section of Analyzer Operating Instructions.
- · Execute recalibration according to the calibration interval required in this package insert.

Quality Control

- When new lot used, check or edit the quality control information.
- Scan the control barcode, choose corresponding quality control information and execute testing. For specific information on ordering quality controls, refer to the
 quality control section of the Analyzer Operating Instructions.

Sample Testing

- After successfully loading the sample, select the sample in interface and edit the assay for the sample to be tested and execute testing. For specific information on
 ordering patient specimens, refer to the sample ordering section of the Analyzer Operating Instructions.
- To ensure proper test performance, strictly adhere to Analyzer Operating Instructions.

Calibration

Traceability: This method has been standardized against the Snibe internal reference standard.

Test of assay specific calibrators allows the detected relative light unit (RLU) values to adjust the master curve.

Recalibration is recommended as follows:

- Whenever a new lot of Reagent or Starter 1+2 is used.
- Every 7 days.

• The analyzer has been serviced.

• Control values lie outside the specified range.

Quality Control

Controls are recommended for the determination of quality control requirements for this assay and should be run in singlicate to monitor the assay performance. Refer to published guidelines for general quality control recommendations, for example Clinical and Laboratory Standards Institute (CLSI) Guideline C24 or other published guidelines ¹².

Quality control is recommended once per day of use, or in accordance with local regulations or accreditation requirements and your laboratory's quality control procedures, quality control could be performed by running the MPO assay:

- Whenever the kit is calibrated.
- Whenever a new lot of Starter 1+2 or Wash Concentrate is used.

Control are only applicable with MAGLUMI and Biolumi system and only used matching with the same top seven LOT numbers of corresponding reagents. For each target value and range refer to the label.

The performance of other controls should be evaluated for compatibility with this assay before they are used. Appropriate value ranges should be established for all quality control materials used.

Control values must lie within the specified range, whenever one of the controls lies outside the specified range, calibration should be repeated and controls retested. If control values lie repeatedly outside the predefined ranges after successful calibration, patient results must not be reported and take the following actions:

- Verify that the materials are not expired.
- Verify that required maintenance was performed.
- Verify that the assay was performed according to the package insert.
- If necessary, contact Snibe or our authorized distributors for assistance.

RESULTS

Calculation

The analyzer automatically calculates the MPO concentration in each sample by means of a calibration curve which is generated by a 2-point calibration master curve procedure. The results are expressed in ng/mL. For further information please refer to the Analyzer Operating Instructions.

Interpretation of Results

The expected ranges for the MPO assay were obtained by testing 326 apparently healthy people in China, and gave the following expected value: ≤94.0 ng/mL (95th percentile).

Results may differ between laboratories due to variations in population and test method. It is recommended that each laboratory establish its own reference interval.

- Results should be used in conjunction with patient's medical history, clinical examination and other findings.
- If the MPO results are inconsistent with clinical evidence, additional testing is needed to confirm the result.
- Specimens from patients who have received preparations of mouse monoclonal antibodies for diagnosis or therapy may contain human anti-mouse antibodies (HAMA). Such specimens may show either falsely elevated or depressed values when tested with assay kits which employ mouse monoclonal antibodies ^{13,14}. In addition, patient samples could contain perinuclear anti-neutrophilic cytoplasmic antibodies (p-ANCA), autoantibodies specific for myeloperoxidase, which could inhibit the immunoreaction in the assay to give falsely depressed results. Additional information may be required for diagnosis.
- Heterophilic antibodies in human serum can react with reagent immunoglobulins, interfering with *in vitro* immunoassays. Patients routinely exposed to animals or animal serum products can be prone to this interference and anomalous values may be observed ¹⁵.
- Bacterial contamination or heat inactivation of the specimens may affect the test results.

SPECIFIC PERFORMANCE CHARACTERISTICS

Representative performance data are provided in this section. Results obtained in individual laboratories may vary.

Precision

Precision was determined using the assay, samples and controls in a protocol (EP05-A3) of the CLSI (Clinical and Laboratory Standards Institute): duplicates at two independent runs per day for 5 days at three different sites using three lots of reagent kits (n = 180). The following results were obtained:

Sampla	Mean (ng/mL)	Within-Run		Between-Run		Reproducibility	
Sample	(n=180)	SD (ng/mL)	%CV	SD (ng/mL)	%CV	SD (ng/mL)	%CV
Plasma Pool 1	97.332	3.397	3.49	1.527	1.57	4.328	4.45
Plasma Pool 2	296.683	7.110	2.40	5.374	1.81	9.490	5.07
Plasma Pool 3	709.400	14.345	2.02	5.123	0.72	23.425	3.30
Control 1	147.893	4.280	2.89	2.121	1.43	6.216	4.20
Control 2	595.361	12.219	2.05	7.016	1.18	18.869	3.17

Linear Range

3.00-1300 ng/mL (defined by the Limit of Quantitation and the maximum of the master curve).

Reportable Interval

2.00-6500 ng/mL (defined by the Limit of Detection and the maximum of the master curvexRecommended Dilution Ratio).

Analytical Sensitivity

Limit of Blank (LoB) =1.00 ng/mL.

Limit of Detection (LoD) =2.00 ng/mL.

Limit of Quantitation (LoQ) = 3.00 ng/mL.

Analytical Specificity

Interference

Interference was determined using the assay, three samples containing different concentrations of analyte were spiked with potential endogenous and exogenous interferents in a protocol (EP7-A2) of the CLSI. The measurement deviation of the interference substance is within ±10%. The following results were obtained:

Interference	No interference up to	Interference	No interference up to
Bilirubin	20 mg/dL	Rheumatoid factor	1500 IU/mL
Hemoglobin	500 mg/dL	ANA	6(S/CO) strong positive
Intralipid	1500 mg/dL	Acetylsalicylic acid	60mg/dL
Acetaminophen	0.025mg/dL	Phenobarbital	10 mg/dL
Nitroglycerin	0.16 µg/mL	Procainamide	10 mg/dL
Simvastatin	32 µg/mL	Biotin	50 ug/ml
HAMA	30 ng/mL	Biotin	30 µg/me

Cross-Reactivity

Cross-reactivity was determined using the assay, three samples containing different concentrations of analyte were spiked with potential cross-reactants in a protocol (EP7-A2) of the CLSI. The measurement deviation of the interference substance is within ±10%. The following results were obtained:

Cross-reactant	No interference up to	Cross-reactant	No interference up to
C-Reactive Protein	550 nmol/L	Lactoperoxidase	800 nmol/L
Immunoglobulin A	400 nmol/L	Troponin I	2000 nmol/L
Thyroid Peroxidase	600 nmol/L	α-1 Antitripsina	1250 nmol/L
Lysozyme	4500 nmol/L	Elastase	2500 nmol/L
Eosinophil peroxidase	922 nmol/L	Lactoferrin	800 nmol/L

High-Dose Hook

No high-dose hook effect was seen for MPO concentrations up to 7000 ng/mL.

Method Comparison

A comparison of the MPO assay with a commercially available immunoassay, gave the following correlations (ng/mL):

Number of samples measured: 116

Passing-Bablok: y=0.9975x-0.2471, T=0.997.

The clinical specimen concentrations were between 7.384 and 1297.080 ng/mL.

REFERENCES

- 1. Abu-Soud H M, Hazen S L. Nitric oxide modulates the catalytic activity of myeloperoxidase [J]. Journal of Biological Chemistry, 2000, 275(8): 5425-5430.
- 2. Arnhold J, Furtmüller P G, Obinger C. Redox properties of myeloperoxidase [J]. Redox report, 2003, 8(4): 179-186.
- Tiruppathi C, Naqvi T, Wu Y, et al. Albumin Mediates the Transcytosis of Myeloperoxidase by Means of Caveolae in Endothelial Cells [J]. Proceedings of the National Academy of Sciences of the United States of America, 2004, 101(20): 7699-7704.
- 4. Klebanoff SJ. Myeloperoxidase: friend and foe [J]. J Leukoc Biol. 2005; 77(5):598-625.
- 5. Nicholls S J, Hazen S L. Myeloperoxidase, modified lipoproteins, and atherogenesis [J]. Journal of Lipid Research, 2009, 50 Suppl: S346-351.
- 6. Nicholls S J, Hazen S L. Myeloperoxidase and cardiovascular disease [J]. Arterioscler Thromb Vasc Biol, 2005, 25(6): 1102-1111.
- 7. Malakar AK, Choudhury D, Halder B, Paul P, Uddin A, Chakraborty S. A review on coronary artery disease, its risk factors, and therapeutics [J]. J Cell Physiol. 2019; 234(10):16812-16823.
- 8. Ndrepepa G, Braun S, Mehilli J, et al. Myeloperoxidase level in patients with stable coronary artery disease and acute coronary syndromes [J]. European Journal of Clinical Investigation, 2008, 38(2): 90-96.
- 9. Zhang, Renliang, Association Between Myeloperoxidase Levels and Risk of Coronary Artery Disease [J]. Jama, 2001, 286(17):2136-2142.
- 10. Roman R M, Camargo P V, Borges F K, et al. Prognostic value of myeloperoxidase in coronary artery disease: comparison of unstable and stable angina patients [J]. Coronary Artery Disease, 2010, 21(3): 129-136.
- 11. Shih J, Datwyler S A, Hsu S C, et al. Effect of collection tube type and preanalytical handling on myeloperoxidase concentrations [J]. Clinical chemistry, 2008, 54(6): 1076-1079.
- 12. CLSI. Statistical Quality Control for Quantitative Measurement Procedures: Principles and Definitions. 4th ed. CLSI guideline C24. Wayne, PA: Clinical and Laboratory Standards Institute; 2016.
- 13. Robert W. Schroff, Kenneth A. Foon, Shannon M. Beatty, et al. Human Anti-Murine Immunoglobulin Responses in Patients Receiving Monoclonal Antibody Therapy [J]. Cancer Research, 1985, 45(2):879-885.
- 14. Primus F J, Kelley E A, Hansen H J, et al. "Sandwich"-type immunoassay of carcinoembryonic antigen in patients receiving murine monoclonal antibodies for diagnosis and therapy [J]. Clinical Chemistry, 1988, 34(2):261-264.

15. Boscato L M, Stuart M C. Heterophilic antibodies: a problem for all immunoassays [J]. Clinical Chemistry, 1988, 34(1):27-33.

SYMBOLS EXPLANATIONS

	Consult instructions for use	AAA	Manufacturer
2°C 8°C	Temperature limit (Store at 2-8 °C)	\Box	Use-by date
Σ	Contains sufficient for <n> tests</n>	*	Keep away from sunlight
<u>††</u>	This way up	EC REP	Authorized representative in the European Community
IVD	In vitro diagnostic medical device	CONTENTS	Kit component
REF	Catalogue number	LOT	Batch code
CE	CE marking		

MAGLUMI® and Biolumi® are trademarks of Snibe. All other product names and trademarks are the property of their respective owners.



Shenzhen New Industries Biomedical Engineering Co., Ltd.

No.23, Jinxiu East Road, Pingshan District, 518122 Shenzhen, P.R. China Tel: +86-755-21536601 Fax:+86-755-28292740



Shanghai International Holding Corp. GmbH (Europe) Eiffestrasse 80, 20537 Hamburg, Germany

Tel: +49-40-2513175 Fax: +49-40-255726